

ing theories of greenhouse warming with Earth's temperature record over the past century and for estimating the response to time-evolving concentrations of greenhouse gases next century. I believe the results I have discussed make a case for diagnosis and intercomparison of oceanic general circulation models, analogous to comparison of atmospheric models⁴, with emphasis on rate of response to external perturbations.

CURT COVEY

Atmospheric and Geophysical Sciences
Division,
Lawrence Livermore National
Laboratory,
Livermore, California 94551, USA

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Rhythms of war

SIR — Schreiber *et al.* proposed¹ that seasonal rhythms influence the opening dates of wars, based on the observed significant correlation between daily photoperiod and the dates of commencement of acts of aggression². They assigned this correlation to increased aggression induced by long photoperiods. Their published correlations were indeed quite remarkable for data from both the Northern and Southern Hemispheres and from the Equator.

However, as with all such comparisons and correlations, an understanding of underlying principles and of the nature of the dataset should be applied to the interpretation of the results. What this high correlation may in fact identify is a relationship between the season and military strategy over the past few thousand years. It seems logical that military strategists should make use of good conditions (including long photoperiod) to move men and equipment, and that they would plan campaigns accordingly. The failure to respond to climate conditions while planning and executing wars has been illustrated by more than one disastrous campaign.

ROBERT BENNETT

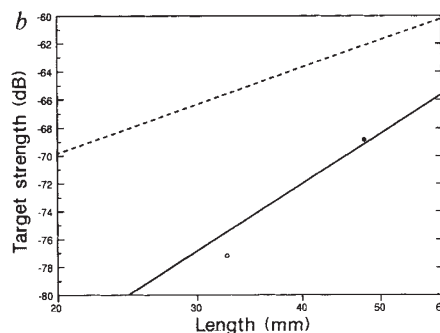
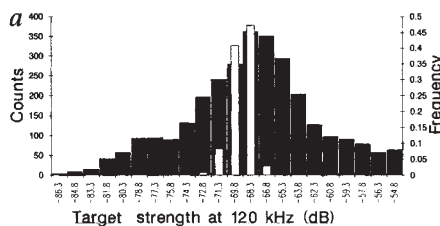
Imperial Cancer Research Fund,
London WC2A 3PX, UK

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Krill abundance

SIR — Antarctic krill (*Euphausia superba*) is the primary food source for many animals in the southern ocean and is also the basis of a large fishery. To manage this resource, krill abundance has been estimated directly with acoustics and indirectly with estimates of predator consumption rates. However, the abundance estimates using acoustics are often an order of magnitude less than those based on predator consumption rates¹. The acoustic method converts echo energy to absolute biomass by assuming that the echo return is the sum of individual scatterers, and by assuming an empirical or modelled acoustic target strength (TS) for individual krill. Everson *et al.*² and Greene *et al.*³ reported new TS measurements of experimentally constrained krill, but to date no corroborating field data have been published. We present new *in situ* TS measurements of krill obtained in March 1991 off Elephant Island, Antarctica.

Foote *et al.*¹ ensouffied live krill aggregations in a cage at 120 kHz. The mean single-animal target strength of krill (lengths 30–39 mm) was inferred from the aggregation backscatter to range from –81 to –74 dB. Past acoustic surveys have typically used TS values



a, TS distribution for 2,957 individual krill detected with a Simrad EK500 echo sounder and a 120-kHz split-beam transducer off Elephant Island, Antarctica in March 1991. Solid bars, *in situ* measurements; open bars, distribution predicted from the sampled distribution of animal lengths. b, TS by length relationship at 120 kHz. Solid line reproduced from ref. 3. Also plotted are the median length and TS from the Foote *et al.* experiment (open circle)¹, the data reported here (solid circle), and the BIOMASS equation (dotted line).

calculated using the equations from the BIOMASS⁴ program, but these equations lead to gross underestimates of krill abundance².

Until recently, a fluid sphere model was thought to characterize adequately the acoustic TS of krill. Wiebe *et al.*⁵ ensouffied several species of live, but tethered, zooplankton at 420 KHz and concluded that sound scatter from elongated animals is better described by a bent cylinder model⁶ and that TS is proportional to the volume of an animal rather than its cross-sectional area. Using these data, Greene *et al.*³ predicted krill TS at several frequencies and over a range of animal lengths. The Foote *et al.* data agree with the TS prediction for a mean animal size of 33 mm. Because the bent cylinder model predicts that TS is much more sensitive to animal length than would be expected with the fluid sphere model, additional measurements of krill TS, particularly at different lengths, could provide strong corroboration of refs 1 and 3.

We present *in-situ* TS measurements of krill (mean length = 47.44 mm, $\sigma = 2.92$), around Elephant Island (*a* in the figure). Zooplankton sound scattering depends upon the morphology of the animal as well as its size, shape and orientation⁵. The spread of the distribution is likely due to the size distribution and variable orientation of the ensouffied krill. Some of the high TS observations may be from multiple krill erroneously identified as individual scatterers. Nonetheless, the distribution is centred on –69dB, within 1dB of the prediction by Greene *et al.* Furthermore, the slope between the Foote *et al.* data and the data presented here is in accordance with that predicted by the bent cylinder model.

The TS of 47.44-mm krill estimated from the BIOMASS equations is –62.3 dB, 6.7 dB above the modal value of the measurements reported here. Thus, for a population composed of 47.44-mm individuals, abundance estimates using the mean TS reported here would be 4.7 times higher than that estimated with the BIOMASS value.

ROGER P. HEWITT

Southwest Fisheries Science Center,
La Jolla, California 92038, USA

DAVID A. DEMER

Scripps Institution of Oceanography,
University of California at San Diego
La Jolla, California 92038, USA

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